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(54) Heat-sealable multilayer polyolefin films

(57) A multilayer polyoletin film of the type suitable or packaging application in which heat seals are formed, and in its preparation the multilayer film comprises a flexible substrate layer formed of a crystalline thermoplastic polymer having an interface surface. A heat-sealable surface layer is bonded to the interface surface of the substrate layer and is formed of a syndiotactic propylene polymer effective to produce a heat seal with itself at a sealing temperature of less than 10°C. The surface layer has a thickness which is less than the thickness of the substrate layer. The heat-seal layer can be formed of syndiotactic polyproplyene polymer of the substrate layer.

lymentraed in the presence of a syndiospecific metallocene catalyst and having a metil flow likes of less than 2 grama*10 minutes. The multilayer film can take the form of a biaxiality-oriented film. In the production of the multilayer film incorporating a substrate layer and a heat-sealable surface layer, a crystalline thermoplastic polymer ie extruded and formed thin a substrate layer film. A second polymer comprising a syndiotactic propylene polymer which is effective to form a heat-sealatibe surface layer is extruded to form a surface layer that is bonded to the interface of the substrate layer at a temperature within the range of 150-280°C. D scripti n

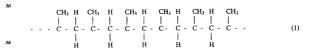
FIELD OF THE INVENTION

5 [0001] This invention reletes to multilayer polyolefin films of the type suitable for peckaging applications involving heat-sealing.

BACKGROUND OF THE INVENTION

(0002) Multileyer polyolefin films incorporete a base or a substrate layer of a stereoreguler crystalline thermoplastic polymer end one or more surface plies which can be formed on one or both sides of the base leyer. Isotactic polypropylene is one of a number of crystelline polymers that can be cheracterized in terms of the stereoregulerity of the polymer chain. Various stereospecific structural relationships denominated primarily in terms of syndiotacticity end isotacticity may be involved in the formation of stereoreguler polymers from various monomers. Stereospecific propegation may be applied in the polymerization of sthylenically unsetureted monomers such as C₂ + sipha olefins, 1-dienes such as 1,3-butadiene, substituted vinyl compounds such as vinyl erometics, e.g. styrene or vinyl chloride, vinyl ethers such as alkyl vinyl ethers, e.g. isobutyl vinyl ether, or even enyl vinyl ethers. Stereospecific polymer propagation is probably of most significance in the production of polypropylene of isotactic or syndiotactic structure.

20 [0003] Isobactic pobypropylene is conventionally used in the production of relatively thin films in which the polypropylene he heated and then activated through dise and subject to baixed inchantion by stressing the film in both elion glitudinal direction (referred to es the mechine direction) and in a transverse or lateral direction sometimes referred to as the "enter" direction. The structure of isotactic polypropylene is characterized in terms of the methyl group statemed to the tertiary carbon atoms of the successive propylene monomer units typing on the same side of the main chain of the polymer. That is, the methyl groups are characterized as being ell above or below the polymer chain. Isotactic polypropylene can be illustrated by the following chemical formula:



Stereoregular polymers, such as isotactic and syndiotactic polypropylene, can be cheracterized in terms of the Fisher o projection formule. Using the Fisher projection formule, the stereochemical sequence of isotectic polypropylene es shown by Formula (2) is described as follows:



Another way of describing the structure is through the use of NMR. Bovey's NMR nomenclature for an isotactic pentad is...mmmm... with each "m" representing a "meso" dyed, or successive methyl groups on the same side of the plane of the polymer chain. As is known in the art, any deviation or inversion in the structure of the chain lowers the degree of isotacticity and crystallinity of the polymer.

[0004] In contrast to the isotactic structure, syndiotectic propylene polymers are those in which the methyl groups attached to the tertiary carbon atoms of successive monomeric units in the polymer chain lie on alternate sides of the plane of the polymer. Using the Fisher projection formula, the structure of syndiotactic polypropylene can be shown as follows:



Syndiotacticity can be characterized in terms of the syndiotactic pentad rrr in which each "r" represents a recemic dyed. Syndiotactic polymers ere semi-crystelline end, like the isotectic polymera, are essentially insoluble in xylene. This crystellinity distinguishes both syndiotactic and isotectic polymers from an atactic polymer, which is non-crystalline and highly soluble in xylene. An atactic polymer exhibits no regular order of repeating unit configurations in the polymer cheln and forms essentially a wew product.

[0005] For many explications the preferred polymer configuration will be a predominantly isotactic or syndiotactic polymer with very little stactic polymer. Catalysts that produce isotactic polyedins are disclosed in U.S. Patant Nos. 4,794,096 and 4,975,403 to Ewen. These patents disclose chiral, stereorigid metellocene catalysts thet polymerize olefins to form isotactic polymers and are especially useful in the polymerization of highly isotactic polymers disclosed. (for example, in the aforementioned U.S. Patant No. 4,794,095, stereorigidity in a metallocene ligand is imparted by means of a structural bridge extending between cyclopentedienty groups. Specifically disclosed in this patent are stereoregular halphum metallocenes that may be characterized by the following formula:

$$R^n(C_5(R^i)_4)_2$$
 HfQp (4)

In Formule (4), (C_S (R')_A) is a cyclopentadienyl or substituted cyclopentadienyl group, R' is independently hydrogen or a hydrocarbyl raddel having 1-20 carbon atoms, and R' is a structural bridge extending between the cyclopentadienyl rings. Q is a halogen or a hydrocarbon radical, such as an alkyl, anyl, alkenyl, alkylenyl, or enylalkyl, heving 1-20 carbon atoms and p is 2.

[0006] Metallocene catalysts, such as those described above, can be used either as so-called "neutral metallocenes" in which case an alumoxene, such as methylelumoxene, is used as a e-o-catelyst, or hey cen be employed as so-called "cationic metallocenes" which incorporate a stable non-coordinating anion and normally of not require the use of an alumoxane. For example, syndiospecific cationic metallocenes are disclosed in U.S. Patent No. 5,243,002 to Razavi. As disclosed there, the metallocene cation is characterized by the cationic metallocene ligand heving sterically dissimiler ring structures that are joined to a positively-charged coordinating transition metal atom. The metallocene cation is essociated with e stable non-coordineting counter-anion. Similar relationships can be esteblished for isospecific metallocenes.

[007] Catalysts employed in the polymertzation of alpha-olefins may be characterized as supported catalysts or unsupported catalysts, sometimes referred to as homogeneous catalysts. Metallocene catalysts are often employed as unsupported or homogeneous catalysts, although, as described below, they also may be employed in supported catalysts components. Traditional supported catalysts are the so-called "conventional" Zlegier-Natta catalysts, such as talenium tetrachloride supported on an active magnesium dichloride es disclosed, for example, in U.S. patent Nos. 4.288, 718 and 4.544,717, both to Mayr of al. A supported catalyst component, as disclosed in the Mayr '718 patent, includes titanium tetrachloride supported on an active anytomic magnesium disclose, such as magnesium disclosed or magnesium disclosed. The supported catalyst component in Mayr '718 is employed in conjunction with a co-catalyst such and an allysiuminum compound, for example, interhylauminum (TEAL). The May '717 patent falcesee selimitar compound that may also incorporate an electron donor compound which may take the form of various amines, phosenes, esters, aldehudes, and alcohols.

[0008] While metallocene catalysts are generally proposed for use as homogeneous catalysts, it is also known in the art to provide supported metallocene catalysts. As disclosed in U.S. patents Nos. 4,701,432 and 4,808,561, both to Welborn, a metallocene cetalyst component may be employed in the form of a supported cetalyst. As described in the Welborn '432 patent, the support may be any support such as taic, an inorganic oxide, or a resinous support material such as a polyoidin. Specific inorganic oxide include silica and alumina, used alone or in combination with other inorganic oxides such as magnesia, zirconia and the like. Non-metallocene transition metal compounds, such as titenium tetrachioride, are also incorporated into the supported catalyst component. The Welborn '561 patent discloses a heterogeneous catalyst that is formed by the reaction of a metallocene and an elumoxane in combination with the support material. A catalyst system embodying both a homogeneous metallocene component and a heterogeneous component, which may be a "conventional" supported Zieder-Natia catalyst. C.a. a supported trainium tetrachioride.

is disclosed in U.S. Patent No. 5,242,876 to Shemshoum et el. Verlous other catalyst systems involving support d metallocene cetelysts are disclosed in U.S. Patent Nos. 5,308,811 to Suge et al and 5,444,134 to Matsumot

[0009] The polymers normelly moley d in the preparation of blaxielly-oriented polypropylene films er usually those prepared through the us of conventionel Ziegler-Nette catalysts of the type disclos d, for exempl, in the afor mentioned patents to Meyr et al. Thus, U.S. Patent No. 5,573,723 to Pelfer et al discloses e process for producing blaxielly-tioned patents to Meyr et al. Thus, U.S. Patent No. 5,573,723 to Pelfer et al discloses e process for producing blaxielly-oriented polypropylene film herving a bese leyer formed of an isotactic polypropylene homopolymer or propylene ethorence opolymers. Other co-polymers of propylene and alpha-oleffins having from 4.8 carbon atoms also mey be employed in the Pelffer process. Thus, the bese leyer mey take the form of e mixture of isotactic polypropylene ethorylene copolymers with resin polymers such as styrene homopolymer having a softening point of about 130-180°C. The surface layer or leyers mey likewise teke the form of e propylene homopolymer or copolymer or the

seme type employed in the base lever. [0010] Processes for the preparetion of blaxlelly-oriented polypropylene films employing polymers produced by the use of isospecific metallocenes involving di- or tri- substituted indenyl groups are disclosed in Cenadian Patent Appilcation No. 2,178,104. Four isotectic polymers disclosed there are besed upon the polymerization of propylene in the presence of heavily substituted bis(indenyi) ligend structures. In each cese, the metallocene used was a sillcon-bridged di-or tri- substituted bis(indenyi) zirconium dichloride. More specifically, the metallocene cetalysts used ere identified in the aforementioned Canadien patent as rac-dimethylsilenediethyl bis(2-methyl-4,6 diisopropyl-1 indenyl) zirconium dichloride, 2 rac-dimethylsilanedlethyl bls(2-methyl-4,5-benzo-1-indenyi) zirconium dichloride, 3 rac-dimethylsilanedlethyl bls(2-methyl-4,5-benzo-1-indenyi) ethyl bis(2-methyl-4-phenyl-1-indenyl) zirconium dichloride, end 4 rac-dimethylsilenediethyl bis(2-ethyl-4-phenyl-1-indenyl) denyl) zirconium dichloride. The verious polymers produced by these metallocenes catalysts are charecterized in terms of moleculer weight, molecular weight distribution, melting point, meltilow index, mean isotectic block length, end isotactic index as defined in terms of mm trieds. The polymers produced had isotectic indices, es thus defined, of ebout 97-98% as contrested with an isotectic index of 93% for a commercial polypropylene compared with a conventional Ziegier-Netta catalyst and molecular weight distributions ranging from about 2.0 to 3.0 as contrasted with a molecular weight distribution of 4.5 for the polypropylene produced by the conventional Zlegler-Nette cetelyst. Similarly, as in the case of the aforementioned patent to Peiffer et el, the Canadian '104 application discloses multileyer films in which the base ply and one or two top piles can be formed of the same or different propylene polymers including propylene homopolymers or copolymers or terpolymers. Where a propylene homopolymer is employed in the top ply, it is described as having a melting point of at least 140°C. Similarly, as in the case of the eforementioned patent to Pelifer et al, the Canadien '104 application discloses multileyer films in which the base ply and one or two top piles can be formed of the same or different propylene polymers including propylene homopolymers or copolymers or terpolymers. Where a propylene homopolymer is employed in the top ply, it is described as heving a melting point of at least 140°C end a melt flow Index of 1 to 20 grams/10 minutes. In the Canedien '104 application a typical film structure, the base ply is cherecterized as providing at least 40% and typically 50-98% of the total film thickness with the outer ply or piles supplying the remainder of the film thickness. Specific overell film thicknesses disclosed in the Canadian '104 application renge from 4 to 100 microns end more specificelly 6 to 30 microns with the base ply specificelly renging from 1.5 to 10 microns and the outer plies from 0.4 tol.5 microns.

SUMMARY OF THE INVENTION

[0011] In accordance with the present invention, there is provided a mutiliayer polyolefin film of the type suitable for packaging application in which heat seals are formed. The mutiliayer film comprises a floxible substrate layer formed of a crystelline thermoplastic polymer heving an interface surface. A heat-sealable surface layer is bonded to the interface of the substrate layer. The surface layer is formed of a syndiotactic propylene polymer which is effective to produce a heat seel with itself et a sealing temperature of less than 110°C. The surface layer has a thickness which is less than the thickness of the substrate layer. Preferably, the substrate leyer has en everage thickness within the range of 5-150 microns, and the surface layer has a thickness which is no more then one-half the thickness of the substrate layer and having e thickness within the range of 0.3-50 microns. Preferably, the heat-seal leyer is formed of syndiotactic polypropylene polymerized in the presence of a syndiospecific metallicene catalyst end heving e met flow Index of less then 2 grems/10 minutes.

Preferably, the multilayer film is e biexielly-oriented film.

1012] In a further espect of the Invention, there is provided a process for the production of a multilayer film incorporating a substrate layer end a heet-seelable surface layer. In cerrying out the invention, e crystelline thermoplastic polymer is extruded end formed into e substrate layer film. A second polymer is employed comprising a syndiotactic polymer which is effective to form a heet-seelable surface layer. The propylene polymer is extruded to firm a heet-seelable surface layer. The propylene polymer is extruded to the interface of the substrate layer et a temperature within the range of 150-280°C.

BRIEF DESCRIPTION OF THE DRAWINGS

[0013] Fig. 1 is a schemetic lilustration in isometric view of a tenter frame system that may be imployed in forming blaxially-oriented multilayer films in eccordance with the presint invention.

- 5 [0014] Fig. 2 is a schematic illustration of et nter freme process incorporeting systems for co-extrusion or xtrusion coeting of surface leyers bonded to e substrate layer to produce multilayer films in accordance with the present invention.
 - [0015] Fig. 3 is a graphical presentation of maximum seal strength versus seal temperature for heat-seal films formed of verticus polymers.
- [0016] Fig. 4 is a graphical illustration of average seal strength versus seel temperature for the polymers illustrated in Fig. 3.
 - [0017] Fig. 5 is a graphical illustration of near term hot seal strength as a function of seal temperature for the verious polymers depicted in Fig. 3.
- [0018] Fig. 6 is a graphical illustration of hot seal strength as a function of seal temperature for the various polymers
 depicted in Fig. 3.

DETAILED DESCRIPTION OF THE INVENTION

[0019] Heet-seelable multilayer polyolefin films, such as used in packaging of food terms and the like, are generally formed by biaxial orientation procedures, and the Invention will be described with respect to biaxially-oriented films. However, it is to be recognized that the invention will find application in other multilayer polyolefin films in which enhanced heat-seal and hot tack characteristics are desireble. Blaxially-oriented films can be cheracterized in terms of certain well-defined characteristics relating to their stereoregular structures and aphysical properties, including multi temperatures and shrinkage characteristics, as well as in relatively/ow-c-efficients of friction and relatively high tensile modull and reletively in permeation rates to oxygen and water. Blaxielly-oriented films of the type incorporating the present invention are formed with a heat-sealable surface layer incorporating a particular syndiotactic propylene polymer as described in greater detail below and by using any suitable oriented film production technique, such as the conventionally-used tenter frame process.

[0020] In general, such blaxially-oriented film production can be carried out by any suitable technique, such es disclosed in the aforementioned Cenadian Patent Application No. 2,178,104 to Peiffer et el. As described in the Pelffer et al epplication, the polymer or polymers used to make the film are melted and then passed through an extruder to e slot die mechanism after which it is passed over a first roller, characterized as a chill roller, which tends to solidify the film. The film is then oriented by stressing it in a longitudinal direction, characterized as the machine direction, and in a transverse direction to arrive at a film which can be characterized in terms of orientetion ratios, sometimes also referred to as stretch ratios, in both longitudinal and transverse directions. The mechine direction orientetion is accomplished through the use of two sequentially disposed rollers, the second or fast roller operating at a speed in relation to the slower roller corresponding to the desired orientation ratio. This may alternatively be accomplished through a series of rollers with Increasing speeds, sometimes with additional Intermediate rollers for temperature control and other functions. After the film has been stressed in the machine direction, it is again cooled and then preheated and passed into a leteral stressing section, for example, a tenter frame mechanism, where it is again stressed, this time in the transverse direction. Orientation in the transverse direction is often followed by an annealing section. Subsequently, the film is then cooled and may be subjected to further treatment, such as corona treatment or flame treatment, as described, for example, in the aforementioned Canadian Patent Application No. 2,178,104 or in U.S. Patent No. 4,029,876 to Beatty, the entire disclosures of which are incorporated herein by reference. The film may also be metallicized as described in U.S. Patent No. 4.692,380 to Reid, the entire disclosure of which is incorporated herein by reference. While corona and flame treatment typically occurs immediately following orientation and prior to the initial roll up, metallicizing is typically performed at a separate time and location.

[0021] Multilayer oriented films comprise a substrate layer, sometimes termed a "base layer" or "core layer," formed of a stereoregular propylene polymer, typically isotactic polypropylene homopolymer, chosen for good stiffness end other physical properties with one or more thinner surface layers used for heat-seeling as well as to provide other properties such as improved silp or berrier qualities, etc. Numerous methods exist for producing multilayer films including coextrusion, extrusion costing, extrusion laminating, or standard lamination techniques.

[0022] Heat-seeling can be accomplished by placing the heat-seeling surface layer in contact with a corresponding layer normally having the seme or similar chemical meke-up as the heat-seeling layer and using a combination of heat and pressure to create a seel bonding the two corresponding layers together. The heat-seel layer will be a surface layer in order to be able to contact and seal with another leyer (or with a different section of itself). After seeling, it is possible that the seeled structure may constitute an internal layer in an even more complex multilayer film or composition of multilayer films.

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[0023] The fileacy of a heat-seal layer can be characterized in terms of the heat-seal strength of the product, the seal Intitiation temporature (SIT) and the so-called "hot text" characteristic, that is, the hot seal strength as measured shortly aft ir formation of this laminated film layer. Typically, the hot seal strength will be measured at 250 milliseconds after formation of this surface layer substrate bond and at following time interval of 500 milliseconds. The seal Initiation temperature is the temperature at which the bonding of the surface layer to the corresponding layer begins to occur. Failure of a heat seal can occur through a number of mechanisms that can be characterized in terms of "peet," when stretch, or "terr failure." Failure of a heat seal due to peel is characterized by a seal peeling apart at the interface. Failure due to web strotch occurs due to a differential strength between the web and the heat-seal. Failure because of tear involves a tearing of the web itself at the seal edge.

[0024] The heat-sealable surface layer is hypically formed by coextrusion of the surface layer polymer with the substrate layer polymer. Co-extrusion can be carried out by simultaneously coinciding the polymer of the heat-seal year and the polymer of the substrate layer through a solted die system to form a film formed of an outer layer of the heatsealing polymer and substrate layer of the core polymer. Additional layers can also be coextruded, either as an additional heat-seal layer on the other surface of the substrate layer, or layers serving other functions, such as barriers, anti-block layers, etc. Alternatively, a heat-seal layer can be extrusion coated later in the film making process. Also, other layers can be added to create a more complex film after or contemporaneous with the formation of the subheat-seal layer to core layer relationship. The advantages of the present invention remain so long as the heat-seal layer is confluence to and bonded with the substrate layer.

[0025] Turning now to Fig. 1, there is shown a schematic illustration of a suitable "Tenter Frame" orientation process that may be employed in producing biaxially-oriented polypropylene film in accordance with the present invention. More particularly and with reference to Fig. 1, a source of molten polymer is supplied from a hopper 10 to an extrader 12 and from there to a slot die 14 which produces a flat, relatively that fin 16 at its output Film 16 is applied over a chill roller 18, and it is cooled to a suitable temperature within the range of about 30°-60°C. The film is drawn off the chill roller 13 to a stretching section 20 to which the machine direction orientation occurs by means of idler rollers 22 and 2 31 hat leads to prehet rollers 25 and 26.

[0026] As the film is drawn off the chili roller 18 and passed over the idler rollers, it is cooled to a temperature of about 30°-60°C. In stretching the film in the machine direction, it is heated by preheat rollers 25 and 26 to an incremental temperature increase of about 60°-100°C and then passed to the slow roller 30 of the longitudinal orienting mechanism. The slow roller may be operated at any suitable speed, usually about 20-40 feet per minute. The fast roller 31 is operated at a suitable speed, typically about 150 feet per minute, to provide a surface speed at the circumference of about 4-7 times that of the slow roller in order to orient the film in the machine direction. As the oriented film is withdrawn from the fast roller, it is passed over roller 33 at room temperature conditions. From here it is passed over tandem idler rollers 35 and 36 to a lateral stretching section 40 where the film is oriented by stretching in the transverse direction. The section 40 includes a preheat section 42 comprising a plurality of tandem heating rollers (not shown) where it is again reheated to a temperature within the range of 130°-180°C. From the preheat section 42 of the tenter frame, the film is passed to a stretching or draw section 44 where it is progressively stretched by means of tenter clips (not shown) which grasp the opposed sides of the film and progressively stretch it laterally until it reaches it maximum lateral dimension. Lateral stretching ratios are typically greater than machine direction stretch ratios and often may range from 5-12 times the original width. Lateral stretching ratios of 8-10 times are usually preferred. The concluding portion of the lateral stretching phase includes an annealing section 46, such as an oven housing, where the film is heated at a temperature within the range of 130°-170°C for a suitable period of time, about 1-10 seconds. The annealing time helps control certain properties, and increased annealing can be used specifically to reduce shrinkage. The biaxially $oriented \ film \ is then \ with drawn \ from \ the \ tenter \ frame \ and \ passed \ over \ a \ chill \ roller \ 48 \ where \ it \ is \ reduced \ to \ a \ temperature$ of less than about 50°C and then applied to take-up spools on a take-up mechanism 50. From the foregoing description, It will be recognized that the initial orientation in the machine direction is carried out at a somewhat lower temperature than the orientation in the lateral dimension. For example, the film exiting the preheat rollers is stretched in the machine direction at a temperature of about 120°C. The flim may be cooled to a temperature of about 50°C and thereafter heated to a temperature of about 160°C before it is subject to the progressive lateral dimension orientation in the tenter

[0027] From the foregoing description it will be recognized that biaxially-oriented film can have a number of properties to its advantage during and after the machine processing steps. A relatively low coefficient friction is desirable, both during the biaxially oriented in procedure and in the end use applications of the ultimately-produced biaxially-oriented film. A relatively high stiffness, as indicated by the tensile modulus in both the machine direction and the transverse direction is usually advantageous. Relatively low permeabilities to gas and water are desirable in many applications, in addition, a high shrinkage factor of the processed film, while undesirable in some cases, can be advantageous in other applications, such as where the film is used in stretch wrapping of food products, electrical components, and the

[0028] Fig. 2 is a schematic diagram illustrating a tenter-frame process carried out with the co-extrusion of one or

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two surface layers with a substrate lay r. The main xiruder 100 is flanked by two supplem nital xirud rs 102 and 104. Through the operation of one of the supplemental extruders 102 or 104, a separate polymer or polym r bi nd may be extruded to be in contact with the primary polymer or polymer bland forming the substrate supplied from the nate solvader 100, if both supplemental xiruders 102 and 104 are us d, th n a sandwich may be created with the primary polymer forming the or or substrate lay r, and the polymers xiruded by the supplem nital extruders 102 and 104 forming surface layers. After extrusion and casting, the multilayer film continues through the machine direction orientation section 106, pre-heating section 108, transverse direction orientation section 110, annealing section 104 coroling section 114, corons treating section 118, and finally the take-up for wind-up) section 118. In an alternative mode of operation, on one or more surface layers may be added in extrusion coating section 120, after machine direction orientation, but before transverse direction orientation. In extrusion coating section 120, after machine direction in the surface is a trivially oriented film emerging from machine direction inentations occur 100, or may be a multilayer film created by co-extrusion by a combination of main extruder 100 and supplemental extruders 102 and or 100 and supplemental extruders 10

[0029] The syndiotactic propylene polymer employed in forming the heal seal layer of the present invention can be characterized by a low seal initiation temperature (SIT) of less than 100°C and effective seal strength characteristics at relatively low seal temperatures of about 110°C or less. The seal initiation temperatures and the low temperature halt-seal strength are substantially less than the corresponding values observed for isotactic polypropylene conventionally used in forming heat-seal layers. In fact, the SIT and heat-seal strength characteristics, together with the hot tack properties of the syndiotactic polypropylene, are generally better for the syndiotactic polypropylene film than for corresponding films produced with ethylene propylene copolymers.

[0030] The syndiotacitic polypropylenes employed in the present invention are produced by the polymerization of propylene in the presence of a syndiospecific metallocene catalyst of the types disclosed in U.S. Patient Nos. 4,892,851 to Ewen et al., 5,225,500 to Ekider et al., and 5,243,002 to Razavi. The syndiospecific metallocenes may be employed as homogeneous catalyst systems, or they may be employed as supported catalyst systems as disclosed, for example, in U.S. Patient No. 5,807,800 to Shamshoum et al. For a further description of suitable syndiospecific metallocene in U.S. Patient No. 5,807,800 to Shamshoum et al. For a further description of suitable syndiospecific metallocene catalyst systems which can be employed in the polymerization of propylene to produce syndiotactic polypropylene, reference is made to the aforementioned patients to Even et al. Elder et al., Razavi, and Shamshoum et al, the entire disclosures of which are incorporated herein by reference.

[0031] As described in greater detail below, the syndiotactic polypropylene employed in forming the heat-seal layer of the present invention is characterized by a melt flow index which is less, usually substantially less, than the various other crystalline polymers or copolymers useful in forming heat seal seyers. As a practical matter, the syndiotactic polyproproplems is characterized by a melt flow index of less than 3 grams/10 minutes and preferably less than 2 grams/10 minutes. The melt flow index is characterized as the melt flow rate as determined in accordance with ASTM Standard 10 minutes. The melt flow index is characterized as the melt flow rate as determined in accordance with ASTM Standard 10 minutes. The melt flow index is characterized as the melt flow rate as determined in accordance with ASTM Standard 10 minutes. The melt flow index is characterized as the melt flow rate as determined in accordance with ASTM Standard 10 minutes.

[0032] The heat-seal layer (or layers) and substrate layer are normally provided in configurations in which the surface layer has a thickness substantially less than the thickness of the substrate layer. For typical packaging applications, the substrate layer will exhibit an average thickness within the range of 5-150 microns. The heat-seal layer will have a thickness of the substrate layer and usually less than 1/3 of the substrate layer. The surface layer than one-half the thickness of the substrate layer and usually less than 1/3 of the substrate layer. The surface layer typically will have a thickness within the range of 0.3-50 microns.

[0033] The substrate layer may be formed of various polymers or polymer blends as described previously, Isotactic polypropylene homopolymers or propylene/ethylene copolymers, typically containing no more than 10 wt. 8 whylene, may be used to form the substrate layer. A preferred substrate layer horporating polypropylene having a very high isotacticity is defined in terms of meso pentads and meso diads but also having irregularities in the polymer structure characterized in terms of 2,1 insertions as contrasted with the predominate 1,2 insertions characteristic of isotactic polypropylene. Thus the polymer chain of the isotactic polypropylene is characterized by intermittent head to head insertions to result in a polymer structure as exemplified below.

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As shown by the polymer structure depicted by Formula (5), the occasional head-to-head insertion resulting from the use of the 2-alkyl substituted indenyl group results in adjacent pendant methyl groups separated by ethylene groups resulting in a polymer structure which b haves, somewhat in the fashion of a random ethylene propylene copolymer and results in e verieble melting point. This results in e polymer which cen be advantageously employed to produce a biaxially-oriented film having good characteristics in terms of strength in both the mechine and transvers directions, low co-efficients for friction, end reletively low permeabilities to where and to oxygen. At the same time, the biaxially-oriented films thus produc d hav satisfactory haze properties, normally less than 1%, and good gloss charact ristics, greater then 90%. This polymer cen be prepared by the polymerization of propylene in the presence of a metellocene catalayst characterized by the formula

in Formule (6), R', R' are each independently e C_1 - C_4 elkyl group or an phenyl group; ind is en indenyl group substituted at the proximal position by the substitutent R_2 and otherwise unsubstituted; Ri is an ethyl, methyl, isopropyl, or tertiary butlyl group; Me is a transition metal selected from the group consisting of ittanium, zirconium, hafnium, and vanadium; and each Ω is independently a hydrocarbyl group or containing 1 to 4 carbon atoms or a halogen.

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[0034] As indicated by Formula (6) above, the sityl bridge cen be substituted with verious substituents in which R' and R' are each independently a methyl group, an ethyl group, a propyl group (including an isopropyl group), and a butlyl group (including a tertiary butlyl or an isobutlyl group). Alternetively, one or both of R', R' can take the place of a phenyl group. Suitable bridge structures are dimethylsityl, diethylsityl, and diphenylshylstructures.

[0035] The RI substituent at the 2 position (the proximal position with regard to the bridge head carbon atom) can be a methyl, ethyl, isopropyl, or tertiary butyl. Preferably, the substituent at the 2 position is a methyl group. As noted previously the indenyl group is otherwise unsubstituded except that it may be a hydrogenated indenyl group. Specificelly, the indenyl ligand can take the form of e 2- methyl indenyl or e 2-methyl tetrahydroindenyl ligand. As will be recognized by those skilled in the art, the ligand structure should be a racemic structure in order to provide the desired enantiomorphic site control mechanism to produce the slotectic polymer configuration.

⁵ [0036] As described previously, the 2.1 insertions produce "mistakes" in the polymer structure which impart the desired non-uniform melting point cheracteristics of the present Invention. The corresponding film is characterized in terms of low water and oxygen permeabilities and low coefficients of friction as described hereinafter. The "mistakes" due to the 2,1 insertions should not however be confused with mistakes resulting in racemic insertions as indicated, for example, by the following polymer structure:

As will be recognized, the structure (7) can be indicated by the pentad mrrm. The "mistekes" corresponding to the heed-to-heed insertion mechanism involved in the present invention are not characterized by or are not necessarily characterized by racemic diads.

[0037] In experimental work certied out with respect to the present invention, the seal strength and hot tack characteristics of tilms formed of spridiotactic pohyprene were valuated against films formed of isolatical proplyine homopolymers end propylene/ethylene copolymers. The syndiotactic pohyprene homopolymers and propylene/ethylene copolymers. The syndiotactic pohypropylene was prepared by the polymertration of propylene in the presence of a syndiospecific bridged metallocene of the type disclosed in U.S. Patent No. 4,892,851 to Even. Exemplary of such metallocene catalyst systems are metallocenes based upon cyclopentedienty fluoreny) ligand structures such as isopropylene (cyclopentedienty fluoreny) zirconium dichioride employed with a occatalyst such as alumoxane. Such syndiotactic polypropylenes can also be prepared inough the use of so-called "cattonic" metallocenes that incorporate a stable non-coordinating anion and do not normally employ the present of an alumoxane. Syndiospedific cationic metallocenes are disclosed, for exemple, in the aforementioned U.S. Patent No. 5,243,002. The syndiotactic polypropylene employed in the experimental work discussed below had a melt flow index of 1.5 g/10 min, and had a minot structure characterized by about 80% syndiotactic pelants (rm.).

[0038] The isotactic polymers employed in the experimental work included both propylene homopolymers and propylene ethylene copolymers prepared by catalysis with isospecific netallocenes as disclosed in the aforementioned perents to Ewen end Ziegier-Nette cetalysts as disclosed in the aforementioned petents to Mayr et al. The characteristics of the metallocene-based proxyl in a homopolymers (identified hergin as MP-1 through MP-4), the metallocenebased propylene ethylene copolym r (Identified herein as MEP-5), the Ziegier-Natta homopolymer (Identified herein as ZP-7), and the Zi gler-Natta propylene/ethylene copolymers (Identified herein as ZEP-8 and ZEP-8) ar set forth in Table I. In Table I the polymers are characterized in terms of melt flow index, MFI (the melt flow rat In grams/10 minut s), the ethylene content (EC) in w.% where applicable, the xylene solubles content (XS) of the polym rs, and, where available, th slotactic floatx I as indicated by the percent of mose pentate).

TABLE I

Polymer	MFL	EC .	nell green	XS.
MP1	5.5		94.0	.3
MP2	13.7		94.1	.4
MP3	9.4		94.0	.5
MP4	3.3		96.3	.2
MEP5	5.0	0.4	95.8	.7
ZEP6	4.9	6.8		9.9
ZP7	8.9			3.2
ZEP8	5.0	6.1		10.0

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[0039] In the experimental work seal strength and hot tack characteristics of the syndiotactic polypropylene and in the above-identified polymers were evaluated on cast films of 50 micross. In carrying out the experimental work, the seal strengths were evaluated on films formed at temperatures starting below the seal initiation temperature to temperatures indicating a plateau in the seal strength characteristic.

[0040] In Fig. 3, the maximum seal strength (SM) in newtons per centimeter is plotted on the ordinate versus the seal temperature T in degrees Centigrade on the absocksar for some of the above-described polymer systems. In Fig. 3, the curves are designated by the reference numerals SP for the syndiotactic polypropylene and by the designations shown in Table 1 for the several listiactic propylene homopolymers or copolymers.

[0041] Fig. 4 shows graphs of average seal strength SA plotted on the ordinate versus the seal temperature in C* plotted on the abscissa. From an examination of Figs. 3 and 4, it can be seen that the syndiotatic polypropylene in and the films formed with the Ziegler-Natta-based copolymers ZEP-8 and ZEP-8 exhibited much lower seal initiation temperatures (SIT) then the Ziegler-Natta homopolymers. Syndiotactic polypropylene exhibited a SIT of about 94°C, 8-7' below that of the copolymer ZEP-8, the highest ethylene content copolymer tested. The Ziegler-Natta copolymer had a somewhat higher seal initiation temperature of about 109°C, and as indicated, the seal temperatures of the remaining polymers were much higher, indicating SIT values of about 120-130°C or even higher. As further indicated by the heal seal strength, the syndiotactic polypropylene develops maximum seal strength over a relatively broad politacui in the low temperature range. While temperatures within the range of 95-115°C can be employed to achieve

effective heal seals, maximum results are achieved within the range of 100-110°C. [0042] The hot tack performance of the syndiciactic polypropylene and the other polymers tested is shown in shown in Figs. 5 and 5, which are grephs of the hot seal strength Sh in newtons per centimeter plotted on the ordinate versus the seal temperature T in °C plotted on the abscless. In each of Figs. 5 and 6, the curves showing the hot teach reacher activities of the polymers are designated by the same reference characters as found in Table 1 and used in Figs. 3 and 4. Fig. 5 indicates the hot seal strength, as measured at 250 milliseconds after bonding and Fig. 6, the hot seal strength at 500 milliseconds after bonding. As can be seen from an examination of Figs. 5 and 6, the syndiciactic polypropylene film demonstrated hot tack performance which is substantially superior to the (metallocene-based and 2Eejler-Natta-based propylene-dethylane copolymers, in the hot tack performance of the remaining polymers, the MIPP homopolymers, as well as the Ziegler-Natta homopolymers, they are Indicated substantially inferior in hot tack performance. In fact, at a seal temperature of 195°C, where the Ziegler-Natta-based propylene-dethylene copolymer begins to show a maximum seal strength greater than the syndiotactic propylene, the syndiotactic polypropylene film still shows far superior hot tack performance as measured in the near term and also the long term.

[0043] Having described specific embodiments of the present invention, it will be understood that modifications thereof may be suggested to those skilled in the art, and it is intended to cover all such modifications as fall within the scope

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of the append d cialms.

Claims

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- 1. In a multilayer polyolefin film of a type suitable for packaging applications, the combination comprising:
 - (a) a flexible substrate layer formed of a crystalline thermoplastic polymer having an interface surface; and (b) a heat sealable surface layer bonded to the interface surface of said substrate layer formed of a syndiotactic propylene polymer and having a thickness which is less than the thickness of said substrate layer, said surface layer being effective to produce a heat seal with itself at a seal temporature less than 110°C.
- 2. The combination of claim 1 wherein said substrate layer is formed of a stereoregular propylene polymer.
- The combination of claim 1 wherein said substrate layer has an average thickness within the range of 5-150 microns
 and said surface layer has a thickness which is no more than one-half the thickness of said substrate layer.
 - The combination of claim 1 wherein sald surface layer is formed of syndiotactic polypropylene produced by the
 polymerization of propylene in the presence of a syndiospecific metallocene catalyst.
 - The combination of claim 4 wherein said syndiotactic polypropylene is characterized by a melt flow index of less than 3 grams/10 minutes.
- The combination of claim 4 wherein sald syndiotectic polypropylene is characterized by a melt flow index of less
 than 2 grams/10 minutes.
 - 7. The combination of claim 4 wherein said substrate layer is formed of an isotactic propylene polymer.
- The combination of claim 7 wherein said substrate layer is formed of an isotactic propylene polymer produced by
 the polymerization of propylene in the presence of an isospecific metallocene catalyst.
 - The combination of claim 1 wherein sald substrate layer is formed of an ethylene propylene copolymer having an ethylene content of no more than 10 wt.%.
- 35 10. The combination of claim 1 wherein said multilayer film is oriented in at least one direction.
 - 11. The combination of claim 1 wherein said multilayer film is biaxially oriented.
- 12. The combination of claim 1 wherein sald multilayer film produces a maximum seal strength upon the heat sealing of sald surface layer with itself at a temperature within the range of 95-110°C of at least 4 newtons/centimeter.
 - 13. The combination of claim 12 wherein said multilayer film produces a maximum seal strength of at least 4 newtons/ centimeter throughout a predominant portion of the range of 100-110°C.
- 45 14. In a process for the production of a multilayer film having a substrate layer and a surface layer, the process comprising:
 - (a) providing a first crystalline thermoplastic polymer;
 - (b) extruding said propylene polymer and forming said polymer into a substrate layer;
 - (c) providing a second polymer comprising a syndiotactic propylene polymer effective to form a surface layer of said multilayer film:
 - (d) extruding said syndiotactic propylene polymer to form a surface layer; and
 - (e) bonding said surface layer to the interface surface of said substrate layer at a temperature within the range of 150°-250°C to form a multilayer film having a surface layer of said syndiotactic propylene polymer which has a thickness that is less than the hickness of said substrate layer.
 - 15. The process of claim 14 wher in said first polym r is an isotactic propylene polymer.

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- 16. The method of claim 14 wherein said second polymer is syndiotactic polypropylene produced by the polymerization of propylene in the presence of a syndiospecific metallocene catalyst.
- 17. The method of claim 16 wherein said syndiotactic polypropylen is characterized by a melt flow index of less than 3 grams/10 minutes.
- 18. The method of cialm 16 wherein said substrate layer film is formed by orienting said substrate layer form in at least one direction and threafter forming said surface layer by extrusion-coating said syndiotactic polypropylene on to said oriented substrate layer film.
- 19. The process of said claim 14 wherein said multilayer film is formed by co-extruding said first and second polymers through a slotted die system to form a multilayer film comprising a substrate layer of said first polymer and a surface layer of said second polymer and thereafter orienting said film in the machine direction followed by orienting the film in the transverse direction to form a blaxially-oriented multilayer film.
- 20. In a process for the production of a multilayer film having a substrate layer and a surface layer, the process comprising:
 - (a) providing a first polymer to form the substrate layer of a multilayer film;

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- (b) providing a second polymer comprising a syndiotactic propylene polymer effective to form a heat-sealable surface layer of said multilayer film; and
- (c) co-extruding said first and second polymers through a slotted die system at a temperature within the range of 150°-250°C to form a film comprising a substrate layer of said first polymer and a surface layer of said second polymer of a thickness which is less than the thickness of said substrate layer.

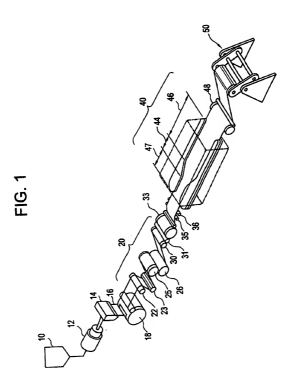


FIG. 2

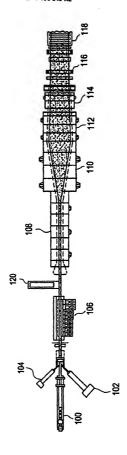
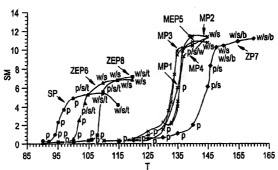
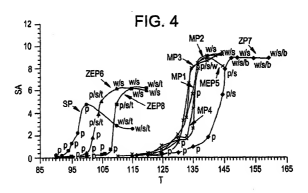


FIG. 3







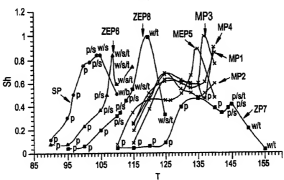


FIG. 6

